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MEMORANDUM

TO:

Mr. Addison Rice

Anderson, Mulholland and Associates

DATE: January 7, 2015

FROM: R. Infante

FILE: 1412152A

RE:

Data Validation

Air samples SDG: 1412152A

SUMMARY

Full validation was performed on the data for several gas samples analyzed for selected volatile organic compounds by method Compendium Method TO-15: Determination Of Volatile Organic Compounds (VOCs) In Air Collected In Specially-Prepared Canisters And Analyzed By Gas Chromatography/Mass Spectrometry (GC/MS), January, 1999. The samples were collected at the Bristol Myer Squib-Building 5 VI facility, Humacao, PR site on December 09, 2014 and submitted to Eurofins Air Toxics, Inc. of Folson, California that analyzed and reported the results under delivery group (SDG) 1412152A.

The sample results were assessed according to USEPA data validation guidance documents in the following order of precedence: Compendium Method TO-15. Determination Of Volatile Organic Compounds (VOCs) In Air Collected In Specially-Prepared Canisters And Analyzed By Gas Chromatography/Mass Spectrometry (GC/MS), January, 1999; Validating Air Samples. Volatile Organic Analysis of Ambient Air in Canisters by Method TO-15, (SOP # HW-31. Revision #4. October, 2006 The QC criteria and data validation actions listed on the data review worksheets are from the primary guidance document, unless otherwise noted.

In general the data is valid as reported and may be used for decision making purposes. The data results are acceptable for use. Several samples exceeded the calibration range and were qualified by the laboratory as estimated values (E).

SAMPLES

The samples included in the review are listed below

Client Sample ID	Lab. Sample ID	Collected Date	Matrix	Analysis
B5IA-8 (2014)	1412152A-01A	12/09/2014	Air	VOCs
B5IA-10 (2014)	1412152A-02A	12/09/2014	Air	VOCs
B5IA-6 (2014)	1412152A-03A	12/09/2014	Air	VOCs
B5IA-7 (2014)	1412152A-04A	12/09/2014	Air	VOCs
B5IA-2 (2014)	1412152A-05A	12/09/2014	Air	VOCs
B5IA-1 (2014)	1412152A-06A	12/09/2014	Air	VOCs
B5IA-AA (2014)	1412152A-07A	12/09/2014	Air	VOCs
B5IA-4 (2014)	1412152A-08A	12/09/2014	Air	VOCs

REVIEW ELEMENTS

Sample data were reviewed for the following parameters, where applicable to the method

- o Agreement of analysis conducted with chain of custody (COC) form
- o Holding time and sample preservation
- Gas chromatography/mass spectrometry (GC/MS) tunes
- o Initial and continuing calibrations
- Method blanks/trip blanks/field blank
- o Canister cleaning certification criteria
- Surrogate spike recovery
- o Internal standard performance and retention times
- o Field duplicate results
- Laboratory control sample/laboratory control sample duplicate (LCS/LCSD) results
- o Quantitation limits and sample results

DISCUSSION

Agreement of Analysis Conducted with COC Request

Sample reports corresponded to the analytical request designated on the chain-of-custody form.

Holding Times and Sample Preservation

Sample preservation was acceptable.

Samples analyzed within method recommended holding time.

GC/MS Tunes

The frequency and abundance of bromofluorobenzene (BFB) tunes were within the QC acceptance criteria. All samples were analyzed within the tuning criteria associated with the method.

Initial and Continuing Calibrations

VOCs (Method TQ-15)

The percent relative standard deviations (%RSDs) and response factors (RFs) of all target analytes were within the QC acceptance criteria in the initial calibration. Correlation coefficients (r²) of target analytes were within the QC acceptance criteria. Ongoing accuracy of the instrument was determined by the analysis of a continuing calibration standard.

Method Blank/Trip Blank/Field Blank

Target analytes were not detected in laboratory method blanks for VOCs.

Summa canister met cleaning certification criteria.

Surrogate Spike Recovery

The surrogate recoveries were within the laboratory QC acceptance limits in all samples analyzed.

Internal Standard Performance

VOCs

Samples were spiked with the method specified internal standard. Internal standard are performance and retention times met the QC acceptance criteria in all sample analyses and calibration standards.

Laboratory/Field Duplicate Results

VOCs

Field/laboratory duplicates were not analyzed as part of this data set. LCS/LCSD results used to assess accuracy; RPD within laboratory control limits.

LCS/LCSD Results

VOCs

LCS/LCSD (blank spike) was analyzed by the laboratory associated with this data package. Recoveries and RPD within laboratory control limits.

Quantitation Limits and Sample Results

Dilutions were required with this data set. Dilution was performed on sample B5IA-1 (2014) due to the presence of high level target species.

Detected results for Toluene, 2-Propanol, and Acetone in samples B5IA-6 (2014), B5IA-7 (2014), B5IA-2 (2014) and B5IA-1 (2014) exceed the instrument calibration range and are considered estimated values.

Detected results for Toluene, 2-Propanol, 4-Methyl-2-pentanone and Acetone in sample B5IA-4 (2014) exceed the instrument calibration range and are considered estimated values.

Calculations were spot checked.

Certification

The following samples 1412152A-01A; 1412152A-02A; 1412152A-03A; 1412152A-04A; 1412152A-05A; 1412152A-06A; 1412152A-07A; and 1412152A-08A were analyzed following standard procedures accepted by regulatory agencies. The quality control requirements met the methods criteria except in the occasions described in this document. The results are valid. Some of the results were qualified.

Méndez

Rafael Infante

Chemist License 1888



Air Toxics

Client Sample ID: B5IA-8 (2014) Lab ID#: 1412152A-01A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name: Dil. Factor:	v121616 1.73	Date of Collection: 12/9/14 11:26:00 AM Date of Analysis: 12/16/14 07:01 PM		
Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (ug/m3)	Amount (ug/m3)
Benzene	0.17	Not Detected	0.55	Not Detected
Ethyl Benzene	0.17	2.7	0.75	12
m,p-Xylene	0.17	10	0.75	44
o-Xylene	0.17	1.0	0.75	4.5
Toluene	0.17	21	0.65	80
2-Propanol	0.86	30	2.1	73
4-Methyl-2-pentanone	0.17	10	0.71	43
Acetone	0.86	45	2.0	100

		Method	
Surrogates	%Recovery	Limits	
1,2-Dichloroethane-d4	111	70-130	
Toluene-d8	106	70-130	
4-Bromofluorobenzene	98	70-130	





Client Sample ID: B5IA-10 (2014) Lab ID#: 1412152A-02A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name: Dil. Factor:	v121617 1.80		Date of Collection: 12/9/14 11:48:00 AM Date of Analysis: 12/16/14 07:43 PM		
Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (ug/m3)	Amount (ug/m3)	
Benzene	0.18	Not Detected	0.58	Not Detected	
Ethyl Benzene	0.18	2.2	0.78	9.5	
m,p-Xylene	0.18	7.7	0.78	34	
o-Xylene	0.18	0.92	0.78	4.0	
Toluene	0.18	17	0.68	64	
2-Propanol	0.90	25	2.2	61	
4-Methyl-2-pentanone	0.18	8.2	0.74	34	
Acetone	0.90	36	2.1	86	

Surrogates	%Recovery	Limits
1,2-Dichloroethane-d4	113	70-130
Toluene-d8	105	70-130
4-Bromofluorobenzene	96	70-130





Client Sample ID: B5IA-6 (2014) Lab ID#: 1412152A-03A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name: Dil. Factor:			Date of Collection: 12/9/14 1:29:00 PM Date of Analysis: 12/16/14 08:41 PM Rpt. Limit Amount (ug/m3) (ug/m3)	
Compound				
Benzene	0.16	0.22	0.50	0.70
Ethyl Benzene	0.16	13	0.69	56
m,p-Xylene	0.16	45	0.69	200
o-Xylene	0.16	4.1	0.69	18
Toluene	0.16	98 E	0.60	370 E
2-Propanol	0.79	160 E	1.9	380 E
4-Methyl-2-pentanone	0.16	50	0.65	210
Acetone	0.79	210 E	1.9	510 E

E = Exceeds instrument calibration range.

Surrogates	%Recovery	Method Limits
1,2-Dichloroethane-d4	102	70-130
Toluene-d8	108	70-130
4-Bromofluorobenzene	102	70-130





Client Sample ID: B5IA-7 (2014) Lab ID#: 1412152A-04A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

Dil. Factor:	V121619 1.78	Date of Collection: 12/9/14 2:20:00 PM Date of Analysis: 12/16/14 09:39 PM		
Compound	Rpt. Limit Amount (ppbv) (ppbv)		Rpt. Limit Am (ug/m3) (ug	
Benzene	0.18	0.23	0.57	0.74
Ethyl Benzene	0.18	18	0.77	81
m,p-Xylene	0.18	68	0.77	290
o-Xylene	0.18	6.2	0.77	27
Toluene	0.18	140 E	0.67	520 E
2-Propanol	0.89	160 E	2.2	400 E
4-Methyl-2-pentanone	0.18	71	0.73	290
Acetone	0.89	240 E	2.1	580 F

E = Exceeds instrument calibration range.

Surrogates	%Recovery	Limits
1,2-Dichloroethane-d4	104	70-130
Toluene-d8	109	70-130
4-Bromofluorobenzene	101	70-130





Client Sample ID: B5IA-2 (2014) Lab ID#: 1412152A-05A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name: Dil. Factor:	v121620 1.74	Date of Collection: 12/9/14 2:4 Date of Analysis: 12/16/14 10:		
Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (ug/m3)	Amount (ug/m3)
Benzene	0.17	0.39	0.56	1.3
Ethyl Benzene	0.17	16	0.76	71
m,p-Xylene	0.17	60	0.76	260
o-Xylene	0.17	5.8	0.76	25
Toluene	0.17	120 E	0.66	450 E
2-Propanol	0.87	150 E	2.1	380 E
4-Methyl-2-pentanone	0.17	60	0.71	250
Acetone	0.87	210 E	2.1	510 E

E = Exceeds instrument calibration range.

Surrogates	%Recovery	Method Limits
1,2-Dichloroethane-d4	100	70-130
Toluene-d8	109	70-130
4-Bromofluorobenzene	104	70-130





Air Toxics

Client Sample ID: B5IA-1 (2014) Lab ID#: 1412152A-06A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name: Dil. Factor:			e of Collection: 12/9/14 2:52:00 PM e of Analysis: 12/16/14 10:54 PM	
Compound			Rpt. Limit (ug/m3)	Amount (ug/m3)
Benzene	0.34	Not Detected	1.1	Not Detected
Ethyl Benzene	0.34	26	1.5	110
m,p-Xylene	0.34	96	1.5	420
o-Xylene	0.34	9.1	1.5	40
Toluene	0.34	190 E	1.3	730 E
2-Propanol	1.7	210 E	4.2	510 E

110

340 E

0.34

1.7

E = Exceeds instrument calibration range.

4-Methyl-2-pentanone

Acetone

Container Type: 6 Liter Summa Canister (100% Certified)

Surrogates	%Recovery	Method Limits
1,2-Dichloroethane-d4	99	70-130
Toluene-d8	108	70-130
4-Bromofluorobenzene	102	70-130



1.4

4.1

440

820 E



4-Methyl-2-pentanone

Acetone

Air Toxics

Client Sample ID: B5IA-AA (2014) Lab ID#: 1412152A-07A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name: Dil. Factor:	v121622 1.67	Date of Collection: 12/9/14 3:08:00 PM Date of Analysis: 12/17/14 05:22 AM			
Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (ug/m3)	Amount (ug/m3)	
Benzene	0.17	Not Detected	0.53	Not Detected	
Ethyl Benzene	0.17	0.20	0.72	0.86	
m,p-Xylene	0.17	0.47	0.72	2.0	
o-Xylene	0.17	Not Detected	0.72	Not Detected	
Toluene	0.17	0.81	0.63	3.0	
2-Propanol	0.84	2.3	2.0	5.6	

0.29

5.1

0.17

0.84

Container Type: 6 Liter Summa Canister (100% Certified)

Surrogates	%Recovery	Method Limits
1,2-Dichloroethane-d4	109	70-130
Toluene-d8	106	70-130
4-Bromofluorobenzene	99	70-130



0.68

2.0

1.2

12



Client Sample ID: B5IA-4 (2014) Lab ID#: 1412152A-08A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

	Rpt. Limit	Amount	Rpt. Limit	Amount	
Dil. Factor: 1.74		Date of Analysis: 12/17/14 06:13 AM			
File Name:	v121623	Dat	e of Collection: 12/9	9/14 3:17:00 PM	

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (ug/m3)	Amount (ug/m3)
Benzene	0.17	0.18	0.56	0.57
Ethyl Benzene	0.17	20	0.76	85
m,p-Xylene	0.17	69	0.76	300
o-Xylene	0.17	6.6	0.76	28
Toluene	0.17	140 E	0.66	520 E
2-Propanol	0.87	160 E	2.1	400 E
4-Methyl-2-pentanone	0.17	71 E	0.71	290 E
Acetone	0.87	240 E	2.1	570 E

E = Exceeds instrument calibration range.

Surrogates	%Recovery	метлоа Limits
1,2-Dichloroethane-d4	103	70-130
Toluene-d8	107	70-130
4-Bromofluorobenzene	102	70-130



Project Manager				n. Housie (800) 48	57-4922	7			age <u> </u>		
Collected by: (Print and Sign)			Project Info:			Turn Around Time:		100	Lab Use Only Pressurized by:		
Company AMAI Email			P.O. #		*	Normal		Date:			
Address 110 Corporale Pk City While Phine	tate WY Zip KO	604	Project # Building S		ing S VI	SVE OR		Dunk Think		urization Gas:	
Phone 914-391-1225 Fax					5 Humancao	l —		1 (53)	1.00	15.	
Lab/I.D. Field Sample I.D. (Location)		,	ate	Time			Canis	ster Pre	N ₂ He ssure/Vacuu	um	
	Can #	of Co	llection	of Collection	Analyses Reques	ted	Initial	Final	PRODUCES AND DESCRIPTION OF THE PARTY OF THE	inal (psi)	
81A-8(2014)	33654	12/	9/14	1126	see notes		28.5	6.5	1	(P81)	
861A-10(2014)	25261		9/14		LI	14. (30+	9.5			
BSIA - 6(2014)	940	12/0	1/14		11	N &c (27.5	2.3			
CHIA BSIA - 7(2014)	33889	12/	4/14	1420	4	*· ·· ··	30	8			
054 BSIA-2 (2014)	3737	12/	3/14	1445	1,	*****	30+	8			
06A BSIA-1 (2014)	12019	12/9	/14.	1452	1,		30	か			
074 BSIA-AA (2014)	5695	12/9	114	1508	1.	*	29.5	5			
OSA B33A - 4 (2014)	25248	12/9	/14	1517	LI		30 r	7"			
Ralinquished by: (signature) Date/Time IRe	,		,					***			
Halinquistred by: (signature) Date/Time Re	ceived by: (signat	ure) D				tone	e, Bei	28 end	, Ellylbe	se	
C - C - C - C - C - C - C - C - C - C -	ceived by: (signat		ate/Time		1010 toluene isoprepy	ryle	ine, M	lethe	inely MIC	3K,	
		u.u, u	, area unit		poterby	AL	charles	Alcol	سا ُن،خ		
Relinquished by: (signature) Date/Time Received by: (signature) D		ate/Time	9	10-15-2	10-15% Methere was Astim						
Shipper Name At Air Bill #					D-1646	·					
Use C. C.		imp (°C	4.1	Condition	Custody Sea	4.25	441 / F	Work C	rder#		
anily redex 77214722 65	347	NA		Cool	Yes No	No	ne)	14	12152	dai:2012	

	Project Number:1412152A Date:12/09/2014
REVIEW OF VOLATILE ORGATHE following guidelines for evaluating volatile organics was actions. This document will assist the reviewer in using prodecision and in better serving the needs of the data users. The USEPA data validation guidance documents in the follow "Compendium Method TO-15. Determination of Volatile Org Specially-Prepared Canisters and Analyzed By Gas Child January, 1999"; USEPA Hazardous Waste Support Branc Analysis of Ambient Air in Canisters by Method TO-15, (SOP QC criteria and data validation actions listed on the data reviewed and the quality control and performance data summater the support of the province of	ANIC PACKAGE ere created to delineate required validation of of pressional judgment to make more informed the sample results were assessed according to ing order of precedence: QC criteria from anic Compounds (VOCs) In Air Collected In romatography/Mass Spectrometry (GC/MS), th. Validating Air Samples. Volatile Organic of #HW-31. Revision #4. October, 2006). The new worksheets are from the primary guidance data package received has been
Lab. Project/SDG No.:1412152ANo. of Samples:8	Sample matrix:Air
Trip blank No.: Field blank No.: Equipment blank No.: Field duplicate No.: X Data Completeness X Holding Times X GC/MS Tuning X Internal Standard Performance X Blanks X Surrogate Recoveries N/A _ Matrix Spike/Matrix Spike Duplicate Overall Comments: _Selected_VOC's_by_method_TO-15	X Laboratory Control SpikesX Field DuplicatesX CalibrationsX Compound IdentificationsX Compound QuantitationX Quantitation Limits
Definition of Qualifiers: J- Estimated results J- Compound not detected R- Rejected data JJ- Estimated nondetect Reviewer: Acul Mauf Date: 01/07/2015	

DATA COMPLETENESS

MISSING INFORMATION	DATE LAB. CONTACTED	DATE RECEIVED
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All criteria were met _X
Criteria were not met
and/or see below

HOLDING TIMES

The objective of this parameter is to ascertain the validity of the results based on the holding time of the sample from time of collection to the time of analysis.

Complete table for all samples and note the analysis and/or preservation not within criteria

SAMPLE ID	DATE SAMPLED	DATE ANALYZED	pН	ACTION
	All samples analyzed w	l rithin the recommended	i method	holding time
···				
			_	
			i i	

<u>Criteria</u>

Aqueous samples – 14 days from sample collection for preserved samples (pH \leq 2, 4°C), no air bubbles.

Aqueous samples – 7 days from sample collection for unpreserved samples, 4°C, no air bubbles.

Soil samples- 7 days from sample collection.

Cooler temperature (Criteria: 4 + 2 °C): N/A – summa canisters

Actions

If the VOCs vial(s) have air bubbles, estimate positive results (J) and reject nondetects (R).

If the % solids of soil samples is 10-50%, estimates positive results (J) and nondetects (UJ)

If the % solid of soil samples is < 10%, estimate positive results (J) and reject nondetects (R).

If holding times are exceeded but < 14 days beyond criteria, estimate positive results (J) and nondetects (UJ).

If holding times are exceeded but < 28 days beyond criteria, estimate positive results (J) and reject nondetects (R).

If holding times are grossly exceeded (> 28 days beyond criteria), reject all results (R).

If samples were not iced or if the ice were melted (> 10°C), estimate positive results (J) and nondetects (UJ).

		Crite	All criteria were metX ia were not met see below
GC/MS TUNING			
The assessment standard tuning (•	determine if the sample instrun	nentation is within the
XThe BFB	performance results were	reviewed and found to be within t	he specified criteria.
XBFB tuni	ng was performed for every	24 hours of sample analysis.	
If no, use profes qualified or reject		ine whether the associated data	should be accepted,
List	the	samples	affected:
If mass calibration	n is in error, all associated	data are rejected.	

All criteria were metX	
Criteria were not met	
and/or see below	

CALIBRATION VERIFICATION

Compliance requirements for satisfactory instrument calibration are established to ensure that the instrument is capable of producing and maintaining acceptable quantitative data.

Date of initial calibration:	_10/06/14	_
Dates of continuing calibration:	12/16/14	
Instrument ID numbers:	MSD-V	
Matrix/Level:	_Air/low	

DATE	LAB	FILE	CRITERIA OUT	COMPOUND	SAMPLES
	ID#		RFs, %RSD, %D, r		AFFECTED
				pecific requirements. Ini	tial calibration retention
times meet	method	specific	requirements.		
		,			

Criteria

All RFs must be > 0.05 regardless of method requirements for SPCC.

All %RSD must be ≤ 15 % regardless of method requirements for CCC.

All %Ds must be \leq 30% regardless of method requirements for CCC.

Method TO-15 does not specify criterion for the curve correlation coefficient (r). A limit for r of \geq 0.995 has therefore been utilized as professional judgment.

Actions

If any compound has an initial RF or a continuing RF of < 0.05, estimate positive results (J) and reject nondetects (R), regardless of method requirements.

If any compound has a %RSD > 15%, estimate positive results (J) and use professional judgment to qualify nondetects.

If any compound has a %RSD > 90%, estimate positive results (J) and reject nondetects (R).

If any compound has a % D > 30%, estimate positive results (J) and reject nondetects (R).

If any compound has a % D > 30%, estimate positive results (J) and nondetects (UJ).

If any compound has a % D > 90%, estimate positive results (J) and reject nondetects (R).

If any compound has r > 0.995, estimate positive results and nondetects.

A separate worksheet should be filled for each initial curve

All criteria were metX
Criteria were not met
and/or see below

V A. BLANK ANALYSIS RESULTS (Sections 1 & 2)

The assessment of the blank analysis results is to determine the existence and magnitude of contamination problems. The criteria for evaluation of blanks apply only to blanks associated with the samples, including trip, equipment, and laboratory blanks. If problems with any blanks exist, all data associated with the case must be carefully evaluated to determine whether or not there is an inherent variability in the data for the case, or if the problem is an isolated occurrence not affecting other data.

List the contamination in the blanks below. High and low levels blanks must be treated separately.

Laboratory blanks

DATE ANALYZED	LAB ID	LEVEL/ MATRIX	COMPOUND	CONCENTRATIO UNITS
All_method	 d_blank_meeth_	 _method_speci	fic_criteria	
Summa_ca	anisters_met_cl	eaning_certifica	ation_criteria	
Field/Equipmen				***************************************
DATE ANALYZED	LAB ID	LEVEL/ MATRIX	COMPOUND	CONCENTRATION UNITS
No_field/trip/equ	uipment_blanks	_analyzed_with	n_this_data_package	
			· · · · · · · · · · · · · · · · · · ·	
				•
				· · · · · · · · · · · · · · · · · · ·

All criteria were metX	
Criteria were not met	
and/or see below	

VB. BLANK ANALYSIS RESULTS (Section 3)

Blank Actions

Action Levels (ALs) should be based upon the highest concentration of contaminant determined in any blank. Do not qualify any blank with another blank. The ALs for samples which have been diluted should be corrected for the sample dilution factor and/or % moisture, where applicable. No positive sample results should be reported unless the concentration of the compound in the samples exceeds the ALs:

ALs = 10x the amount of common contaminants (methylene chloride, acetone, 2-butanone, and toluene)

ALs = 5x for any other compounds

Specific actions are as follows:

If the concentration is < sample quantitation limit (SQL) and \le AL, report the compound as not detected (U) at the SQL.

If the concentration is \geq SQL but \leq AL, report the compound as not detected (U) at the reported concentration.

If the concentration is \geq SQL and > AL, report the concentration unqualified.

Notes:

High and low level blanks must be treated separately

Compounds qualified "U" for blank contamination are still considered "hits" when qualifying for calibration criteria.

CONTAMINATION SOURCE/LEVEL	COMPOUND	CONC/UNITS	AL/UNITS	SQL	AFFECTED SAMPLES
		-			
					1/21 ²
				·	
			-		
() () () () () () () () () ()					

All criteria were met _X
Criteria were not met
and/or see below

SURROGATE SPIKE RECOVERIES

d4

Laboratory performance of individual samples is established by evaluation of surrogate spike recoveries. All samples are spiked with surrogate compounds prior to sample analysis. The accuracy of the analysis is measured by the surrogate percent recovery. Since the effects of the sample matrix are frequently outside the control of the laboratory and may present relatively unique problems, the validation of data is frequently subjective and demands analytical experience and professional judgment.

List the percent recoveries (%Rs) which do not meet the criteria for surrogate recovery. Matrix: solid/aqueous

Δ2	PI	F	ID

SURROGATE COMPOUND

ACTION

1,2-DICHLOROETHANE-

Toluene-

4-BFB

d8

_Surrogate_recoveries_within_laboratory_control_limits						
		-				
QC Limits* (Air)						
LL_to_UL70to_130	_70to_13070to_130					

- * QC limits are laboratory in-house performance criteria, LL = lower limit, UL = upper limit.
- * If QC limits are not available, use limits of 80 120 % for aqueous and 70 130 % for solid samples.

Actions:

QUALITY	%R < 10%	%R = 10% - LL	%R > UL
Positive results	J	J	J
Nondetects results	R	UJ	Accept

Surrogate action should be applied:

If one or more surrogate in the VOC fraction is out of specification, but has a recovery of > 10%.

If any one surrogate in a fraction shows < 10 % recovery.

All criteria were met
Criteria were not met
and/or see belowN/A

VII. A MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD)

This data is generated to determine long term precision and accuracy in the analytical method for various matrices. This data alone cannot be used to evaluate the precision and accuracy of individual samples. If any % R in the MS or MSD falls outside the designated range, the reviewer should determine if there are matrix effects, i.e. LCS data are within the QC limits but MS/MSD data are outside QC limit.

1. MS/MSD Recoveries and Precision Criteria

The laboratory should use one MS and a duplicate analysis of an unspiked field sample if target analytes are expected in the sample. If target analytes are not expected, MS/MSD should be analyzed.

	PD of the compounds			t the criteria. /Level:	<u>.</u>
MS OR MSD	COMPOUND	% R	RPD	QC LIMITS	ACTION
	_are_not_required_as	•		TO-15;_blank_sp	ike_used_to_assess
	ts are laboratory in-ho nits are not available.	•		•	r limit, UL = upper limit.

Actions:

QUALITY	%R < LL	%R > UL
Positive results	J	J
Nondetects results	R	Accept

MS/MSD criteria apply only to the unspiked sample, its dilutions, and the associated MS/MSD samples:

If the % R for the affected compounds were < LL (or 70 %), qualify positive results (J) and nondetects (UJ).

If the % R for the affected compounds were > UL (or 130 %), only qualify positive results (J).

If 25 % or more of all MS/MSD %R were < LL (or 70 %) or if two or more MS/MSD %Rs were < 10%, qualify all positive results (J) and reject nondetects (R).

A separate worksheet should be used for each MS/MSD pair.

All criteria were met _____ Criteria were not met and/or see below __N/A__

VII. B MATRIX SPIKE/MATRIX SPIKE DUPLICATE

MS/MSD - Unspiked Compounds

It should be noted that Method TO-15 does not specify a MS/MSD criteria for the unspiked compounds in the sample. A %RSD of < 50% has therefore been utilized as professional judgment.

If all target analytes were spiked in the MS/MSD, this review element is not applicable.

List the %RSD of the compounds which do not meet the criteria.

Sample ID:	pie ID: Matrix/Level/Unit:				
COMPOUND	SAMPLE CONC.	MS CONC.	MSD CONC.	% RSD	ACTION
	<u> </u>				
				-	
		4.504-24			

Actions:

^{*} If the % RSD > 50, qualify the positive result in the unspiked samples as estimated (J).

^{*} If the % RSD is not calculated (NC) due to nondetected value, use professional judgment to qualify the data.

All criteria were metX	
Criteria were not met	
and/or see below	

VIII. LABORATORY CONTROL SAMPLE (LCS) ANALYSIS

This data is generated to determine accuracy of the analytical method for various matrices.

1. LCS Recoveries Criteria

Where LCS spiked with the same analyte at the same concentrations as the MS/MSD? Yes or No. If no make note in data review memo.

List the %R of compounds which do not meet the criteria

	LCS ID	COMPOUND	% R	QC LIMIT
LCS/LCS	D_(Blank_spike	e)_analyzed_in_this_data_ı	oackage,_recoveries_a	and_RPD
witnin_lai	boratory_contro			

- QC limits are laboratory in-house performance criteria, LL = lower limit, UL = upper
- If QC limits are not available, use limits of 70 130 %.

Actions:

QUALITY	%R < LL	%R > UL
Positive results	J	J
Nondetects results	R	Accept

All analytes in the associated sample results are qualified for the following criteria.

If 25 % of the LCS recoveries were < LL (or 70 %), qualify all positive results (i) and reject nondetects (R).

If two or more LCS were below 10 %, qualify all positive results as (J) and reject nondetects (R).

2. Frequency Criteria:

Where LCS analyzed at the required frequency and for each matrix? Yes or No. If no, the data may be affected. Use professional judgment to determine the severity of the effect and qualify data accordingly. Discuss any actions below and list the samples affected.

		All criteria were metX Criteria were not met and/or see below
IX.	LABORATORY DUPLICATE PRECISION	
	Sample IDs:	Matrix:

Field duplicates samples may be taken and analyzed as an indication of overall precision. These analyses measure both field and lab precision; therefore, the results may have more variability than laboratory duplicates which only laboratory performance. It is also expected that soil duplicate results will have a greater variance than water matrices due to difficulties associated with collecting identical field duplicate samples.

The project QAPP should be reviewed for project-specific information.

Suggested criteria: RPD ± 25% for air samples. If both samples and duplicate are <5 SQL, the RPD criteria is doubled.

COMPOUND	SQL	SAMPLE CONC.	DUPLICATE CONC.	RPD	ACTION
No field/laboratory of			s data package.		CSD used to assess accuracy,

Actions:

Qualify as estimated positive results (J) and nondetects (UJ) for the compound that exceeded the above criteria. For organics, only the sample and duplicate will be qualified.

If an RPD cannot be calculated because one or both of the sample results is not detected, the following actions apply:

If one sample result is not detected and the other is greater than 5x the SQL qualify (J/UJ).

If one sample value is not detected and the other is greater than 5x the SQL and the SQLs for the sample and duplicate are significantly different, use professional judgment to determine if qualification is appropriate.

If one sample value is not detected and the other is less than 5x, use professional judgment to determine if qualification is appropriate.

If both sample and duplicate results are not detected, no action is needed.

All criteria were metX	
Criteria were not met	
and/or see below	

X. INTERNAL STANDARD PERFORMANCE

The assessment of the internal standard (IS) parameter is used to assist the data reviewer in determining the condition of the analytical instrumentation.

List the internal standard area of samples which do not meet the criteria.

- * Area of +40% or -40% of the IS area in the associated calibration standard.
- * Retention time (RT) within ± 0.06 seconds of the IS area in the associated calibration standard.

DAIE	SAMPLE ID	IS OUT	IS AREA	ACCEPTABLE RANGE	ACTION
	tandard_area_and_reration_standards			_control_limits_for_	_both_samples
 					
				· · · · · · · · · · · · · · · · · · ·	
Actions:					

1. IS actions should be applied to the compound quantitated with the out-of-control ISs

QUALITY	IS AREA < -40%	IS AREA > + 40%
Positive results	J	J
Nondetected results	R	ACCEPT

If a IS retention time varies more than 0.330 seconds, the chromatographic profile for that sample must be examined to determine if any false positive or negative exists. For shifts of a large magnitude, the reviewer may consider partial or total rejection of the data for the sample fraction.

All criteria were met _X
Criteria were not met
and/or see below

XII. SAMPLE QUANTITATION

The sample quantitation evaluation is to verify laboratory quantitation results. In the space below, please show a minimum of one sample calculation:

LCS 12/16/14

Acetone

RF = 1.18134

[] = (141344)(5.0)/(108561)(1.18134)

= 5.51 ppbv OK

All criteria were metX
Criteria were not met
and/or see below

- XII. QUANTITATION LIMITS
- A. Dilution performed

SAMPLE ID	DILUTION FACTOR	REASONS FOR DILUTION
1412152A-06A	3.42	High level of target species

B.	Percent Se	olids
D.	L'EIREIII O	אווע

List samples which have ≤ 50 % solids				

Actions:

If the % solids of a soil sample is 10-50%, estimate positive results (J) and nondetects (UJ)

If the % solids of a soil sample is < 10%, estimate positive results (J) and reject nondetects (R) $^{\parallel}$